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Floating-Point L^2 -Approximations

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Abstract: Computing good polynomial approximations to usual functions is an important topic for the computer evaluation of those functions. These approximations can be good under several criteria, the most desirable being probably that the relative error is as small as possible in the L^∞ sense, i.e. everywhere on the interval under study.

In the present paper, we investigate a simpler criterion, the L^2 case. Though finding a best polynomial L^2 -approximation with real coefficients is quite easy, we show that if the coefficients are restricted to be floating point numbers to some precision, the problem becomes a general instance of the CVP problem, and hence is NP-hard.

We investigate the practical behaviour of exact and approximate algorithms for this problem. The conclusion is that it is possible in a short amount of time to obtain a relative or absolute best L^2 -approximation. The main applications are for large dimension, as a preliminary step of finding L^∞ -approximations and for functions with large variations, for which relative best approximation is by far more interesting than absolute.

Key-words: Floating-point arithmetic, efficient approximation, L^2 norm, L^∞ norm, lattice basis reduction, closest vector problem.

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Approximation flottante au sens L^2

Résumé : Calculer de bons polynômes d'approximation pour les fonctions usuelles est un sujet important, par exemple pour obtenir des méthodes performantes d'évaluation de ces fonctions. La qualité desdites approximations peut être mesurée à l'aune de plusieurs critères. Le critère le plus intéressant pour les applications est probablement la minimisation de l'erreur relative au sens L^∞ , c'est-à-dire uniformément sur l'intervalle considéré.

Dans ce travail, nous étudions un critère plus simple, à savoir la minimisation au sens L^2 . Bien que la recherche de l'optimum à coefficients réels soit une tâche aisée, nous montrons que, si les coefficients sont des nombres flottants de précision fixée, le problème devient une instance générale du problème CVP, et est donc NP-difficile.

Nous étudions néanmoins le comportement pratique des algorithmes exacts et approchés pour ce problème. Notre conclusion est qu'il est possible, de façon efficace, d'obtenir la meilleure approximation au sens L^2 , absolu ou relatif. Les principales applications sont la recherche d'approximants en grande dimension, l'utilisation comme étape préliminaire pour trouver des approximations L^∞ , ou pour les fonctions avec de grandes variations, pour lesquelles l'approximation relative est de loin préférable.

Mots-clés : Arithmétique flottante, approximation efficace, norme L^2 , norme L^∞ , réduction de réseaux, recherche de vecteur le plus proche.

1 Introduction

Computing elementary or special functions is a central topic in modern computer arithmetic. One common way of doing this is, given a continuous real-valued function f , to split the domain of f in sufficiently many small intervals; and then, over each of those intervals I , to compute a polynomial P_I such that the distance between P_I and f over I is less than the accuracy expected. That distance is usually measured with the supremum norm (or L^∞ norm or absolute error)

$$\|P_I - f\|_\infty = \sup_{x \in I} |P_I(x) - f(x)|,$$

or the relative error

$$\|P_I - f\|_{\text{rel}} = \sup_{x \in I} \frac{1}{|f(x)|} |P_I(x) - f(x)|.$$

The classical theorem of Stone-Weierstraß (see eg. [5]) tells us that, for any $\varepsilon > 0$, we can find a polynomial such that $\max_{x \in I} \|f(x) - P(x)\|_\infty \leq \varepsilon$, but it gives no way of computing it. Remes' algorithm [17] gives an efficient way of computing this polynomial; however, this best approximant will not have machine number coefficients, except in very special cases. Of course, one can choose to round the coefficients of the polynomial computed by Remes' algorithm so that they fit the imposed finite-precision arithmetic format but this is not at all optimal: we are interested in getting the best accuracy possible with an approximation having the least degree and the smallest number of bits for representing the coefficients possible and this naive rounding process may give very poor results in that direction. A recent work [4] has made progresses towards the computation of best approximations *with machine number as coefficients*.

In this paper, we propose to investigate approximation by polynomials with machine-number as coefficients, but in the L^2 sense, ie. we are trying to minimize an expression

$$\|f - P_I\|_2 = \left(\int_I (f - P_I)^2 d\mu \right)^{1/2}$$

for some positive measure $d\mu$ over I . Not only interesting in itself, a motivation for studying it is that very good L^2 -approximations may constitute good approximations with respect to L^∞ norm or relative error. This is very useful since in several applications, one is not interested in getting the best approximation but rather a good enough one and also since a better estimate beforehand of the optimal error may lead to a significant speedup of the method presented in [4].

This L^2 context is a simpler situation since it comes with a lot of mathematical structure, and allows thus to devise simpler algorithms. In that case, finding the best polynomial of degree $\leq n$ with real coefficients is simply a matter of projecting the function f over the vector space generated by monomials $1, x, \dots, x^n$. This is discussed in Section 2, in a slightly more general setting which might be of use in some applications. Then, in Section 3, we introduce the problem we address in that article. In Section 4, we shall prove that finding the best polynomial (or, more generally, approximation over any family of L^2 -functions) with machine numbers as coefficients is a (general) instance of the so-called CVP problem. We shall review briefly algorithms to solve CVP and give practical results in Section 5. Finally, we shall give some examples and compare the L^2/L^∞ approaches in Section 6, before concluding our paper.

2 L^2 -approximation

Let I be an interval of \mathbb{R} , and $d\mu$ a positive measure on I . We define the classical scalar product on $H = L^2(I, d\mu)$ as

$$(f|g) = \int_I f(t)g(t)d\mu.$$

Notice that $d\mu$ needs not be the Lebesgue measure.

In particular, for any nonnegative weight function $w(x)$ we can consider the measure $w(x)dx$. This allows one to “fine-tune” the accuracy of the approximation in some parts of the interval. For instance if $I = [-a, a]$, the classical kernel $w(x) = (a^2 - x^2)^{1/2}$ will make the approximation sharper in the center of the interval, and more sloppy close to the bounds.

Let now $(e_i)_{0 \leq i \leq n}$ be a linearly independent family of elements of H , and put $E = \oplus_{i=0}^n \mathbb{R}e_i$. The following Lemma is elementary.

Lemma 1 *Let $f \in H$. The element $p_E(f)$ of E which is closest to f in the L^2 sense has coordinates over the basis (e_i) given by $G^{-1}V$, where G is the Gram matrix $((e_i|e_j))_{0 \leq i, j \leq n}$ and V the vector $((f|e_i))_{0 \leq i \leq n}$.*

In this classical setting, the problem under consideration is thus easy; in the sense that it is reduced to $O(n^2)$ computations of integrals and one linear system solution.

In the sequel, we shall denote by $d_E(f) := d(f, E)$ the L^2 distance from f to the space E i.e. $(\int_I (f - p_E(f))^2 d\mu)^{1/2}$. This is a lower bound for the L^2 distance of f to any element of E .

Remark 1 *The “abstract” point of view developed above has the interest of allowing applications which are not restricted to finding best absolute polynomial approximations. This classical setting is obtained by taking $e_i = x^i$, and $d\mu = dx$ the Lebesgue measure, and the reader should probably keep this in mind as a roadmap.*

However, dealing with the problem in generality allows one to find approximation by trigonometric polynomials by taking $e_i = \cos(ix)$, or to find polynomial approximation with constraints on the values of some coefficients by taking $e_i = x^{k_i}$ where $(k_i)_{0 \leq i \leq n}$ is a finite strictly increasing sequence of natural integers, or to find best polynomial approximations to a function f , over an interval where f has no zero over I by taking $d\mu = dx/f(x)^2$, since then

$$\int_I (P(x) - f(x))^2 d\mu = \int_I \left(\frac{P(x)}{f(x)} - 1 \right)^2 dx.$$

These generalizations are obtained with the same algorithm; one does just need to compute slightly different integrals. Note that we may then derive good polynomial or sum of cosines approximations with respect to the L^∞ absolute or relative errors.

3 Floating-point approximations

In the context of floating-point computations, however, approximations such as $p_E(f)$ have a major drawback: their coefficients are (exact) real numbers. A simple idea is to round them

to the nearest machine-number. However, this can easily make the quality of the approximant much worse. In this paper we are interested in finding the best approximation with “floating-point numbers” as coefficients.

We model our problem FP-*appr* in the following way:

FP-*appr* $((e_i)_{0 \leq i \leq n}, f, d\mu, I)$. Given an interval I , a positive measure $d\mu$ over I , functions $(e_i)_{0 \leq i \leq n} \in L^2(I, d\mu)$, a function $f \in L^2(I, d\mu)$, find a vector $P = (p_i)_{0 \leq i \leq n} \in \mathbb{Z}^{n+1}$ which minimizes

$$\left\| \sum_{i=0}^n p_i e_i(x) - f(x) \right\|_2. \quad (1)$$

Note that it is easily proved that such a P exists. To see why this amounts to approximate by floating-point numbers, refer to the case where $e_i(x) = x^i 2^{-m_i}$ for some integers m_i ; in that case, we get an approximation by polynomials with precision m_i floating-point numbers as coefficients (actually not exactly approximation by floating point numbers, since the p_i are not bounded in our problem).

4 Reduction to CVP

The problem described above can be restated in terms of lattices. Let L be the discrete subgroup of H generated by the functions e_i , i.e. $L = \{\sum_{i=0}^n u_i \cdot e_i, u_i \in \mathbb{Z}^{d+1}\}$. Then L , with the restriction of the scalar product is a lattice of E of maximal rank, and the problem under consideration amounts to find the vector of L which is the closest to $p_E(f)$. We are thus facing an instance of the classical CVP problem (in its “Gram form”), which we now state in two different forms:

CVP-*Gram* (G, V) . Given an $n \times n$ symmetric definite positive matrix G and a vector $V \in \mathbb{R}^n$, find the vector $X \in \mathbb{Z}^n$ which makes $(X - G^{-1}V)^t G (X - G^{-1}V)$ minimal.

CVP (M, v) . Given an $n \times n$ matrix M and a vector $v \in \mathbb{R}^n$, find the vector $x \in \mathbb{Z}^n$ such that $|Mx - v|_2$ is minimal, where $|(y_i)_{1 \leq i \leq n}|_2 = (\sum_{i=1}^n y_i^2)^{1/2}$.

The second form easily reduces to the first one, by taking $G = M^t M$ and $V = Gv$. Conversely, the first form can be reduced, at least numerically, to the second, by computing a Cholesky decomposition (a “square root”) of the matrix G .

In fact, we now prove that our problem is indeed a general CVP-*Gram*, which means that any CVP instance can be reduced to it. We start by giving a precise formulation of the problem we are studying. The formulation given below is actually a bit less general than above (we restrict the e_i and f to be polynomials and assume that the measure $d\mu$ is given by its first $2n$ moments), in order to deal with objects that have a finite representation.

Proposition 1 *Let (u_0, \dots, u_n) be $n+1$ linearly independent vectors in \mathbb{R}^{n+1} , L the lattice they generate, and $v \in \mathbb{R}^{n+1}$ a vector. There exist a sequence of polynomials $(P_i)_{0 \leq i \leq n+1}$ with rational integer coefficients, a sequence $(\mu_i)_{0 \leq i \leq 2n}$ of rational numbers such that $\text{CVP-GrAm}(G, V)$ reduces to $\text{FP-appr}((P_i)_{0 \leq i \leq n}, P_{n+1}, d\mu, \mathbb{R})$ where $d\mu$ is a positive measure over \mathbb{R} with $(\mu_i)_{0 \leq i \leq 2n}$ as its first $2n+1$ moments. Further, all those polynomials can be computed in polynomial time.*

Proof. Up to a (rational) change of basis, we can assume that the matrix $(g_{ij})_{0 \leq i, j \leq n}$ is diagonal. This is achieved in practice by Gram-Schmidt orthogonalization, which can be performed (over \mathbb{Q}) in polynomial time. We are thus looking for orthogonal polynomials Q_i for a measure, with given norm $\sqrt{g_{ii}}$. We start with the polynomials and build the moments of the corresponding measure.

First, put $g_{kk} = 1$ for $n+1 \leq k \leq 2n$. Put $\lambda_0 = g_{00}$ and $\lambda_i = g_{ii}/g_{i-1, i-1}$ for $i = 1, \dots, 2n$, and define Q_i by the recurrence relation

$$Q_0 = 1, Q_1 = x, Q_{i+1} = xQ_i - \lambda_i Q_{i-1} \text{ for } i = 1, \dots, 2n.$$

Put $\mu_0 = \lambda_0$. Then, from

$$\int_{\mathbb{R}} Q_0 Q_1 d\mu = 0 = \int_{\mathbb{R}} x d\mu,$$

we deduce that $\mu_1 = 0$.

More generally,

$$0 = \int_{\mathbb{R}} Q_i Q_0 d\mu = \int_{\mathbb{R}} x Q_{i-1} d\mu - \lambda_i \int_{\mathbb{R}} Q_{i-2} d\mu,$$

which, by expanding the polynomials Q_i , allows to find μ_i in terms of the previous μ_j , for all $i = 1, \dots, 2n$. Then, for $k \leq l-1$ one can rewrite $x^k Q_l(x)$ as a linear combination of Q_{l-k}, \dots, Q_{l+k} , hence $\int_{\mathbb{R}} x^k Q_l(x) d\mu = 0$, which shows that Q_l is indeed orthogonal to Q_0, \dots, Q_{l-1} .

Since the g_{ii} are positive, the bilinear form induced by μ on the polynomials of degree $\leq 2n$ is positive definite, so that Hamburger's theorem [9, 10, 11] applies and there exists a positive measure μ over \mathbb{R} with first moments μ_i .

Finally, for $1 \leq i \leq n$, since Q_i is orthogonal to the Q_j for $j < i$, it is orthogonal to any polynomial of degree $< i$. Hence

$$\begin{aligned} \int_{\mathbb{R}} Q_i^2(x) d\mu &= \int_{\mathbb{R}} x^i Q_i(x) d\mu \\ &= \int_{\mathbb{R}} x^{i-1} Q_{i+1}(x) d\mu + \lambda_i \int_{\mathbb{R}} x^{i-1} Q_{i-1}(x) d\mu \\ &= \lambda_i \int_{\mathbb{R}} Q_{i-1}^2(x) d\mu, \end{aligned}$$

which shows, by induction, that $\int_{\mathbb{R}} Q_i^2(x) d\mu = g_{ii}$. \square

This shows, in view of [20], that FP-appr is NP-hard; further, as a corollary from [13], we obtain

Corollary 1 *The problem FP-appr is NP-hard, even if we allow an approximation factor up to $n^{O(1/\log \log n)}$.*

In practice, fortunately, the situation is by far better. First, the problem is polynomial when the dimension is fixed. Second, the exponential algorithms behave quite well in our situation, where we are looking for approximations of small enough dimension (in any case, less than, say, 50).

5 A short review of algorithms for CVP

The previous section shows us that we need to study the behaviour of CVP algorithms in the context of our problem. Though there is already abundant theoretical and practical literature about the CVP (see e.g. [1]), we include this section for the sake of completeness, but only give heuristic descriptions of the algorithms.

In a nutshell, algorithms for CVP can be decomposed into two distinct steps for which various solutions exist. The first step is a preprocessing step, which computes a more or less strongly reduced basis of the lattice under study. The second one actually solves the CVP. Usually, the stronger the reduction, the more accurate or efficient is the second step.

5.1 Short review of CVP algorithms

5.1.1 A naive method

Let us start with an extremely naive method, which should probably not be used but serves as a good introduction. Lemma 1 tells us that the coordinates of $p_E(f)$ in the basis e_i are given by $G^{-1}V$. A simple idea is thus to compute $G^{-1}V$ and to round to the nearest integer the values that we have obtained in order to get a vector with integer coordinates in the basis (e_i) .

Lemma 2 *This method is optimal if the basis (e_i) is orthogonal.*

Proof. Write $(f_i) = G^{-1}V$. Then, since the basis is orthogonal, for any $x \in \mathbb{Z}^n$, the L^2 distance of $\sum_{i=0}^n x_i e_i$ to $p_E(f) = \sum_{i=0}^n f_i e_i$ is given by

$$\left(\sum_{i=0}^n (x_i - f_i)^2 \|e_i\|_2^2 \right)^{1/2},$$

which is minimal for $x_i = \lfloor f_i \rfloor$. □

5.1.2 Babai's nearest plane algorithm

Babai's method [2] is a refinement of the previous method. In a very sketchy way, it can be described as follows: rounding each coefficient $M^{-1}v$ in the naive method induces an error. Thus, a better way is to round each coefficient one after the other, and to reintroduce the error coming from one rounding before performing the next.

Algorithm:Babai**Data:** An $(n + 1) \times (n + 1)$ Gram matrix G ; an $(n + 1)$ -vector V **Result:** An approximation of $\text{CVP-Gram}(G, V)$ **begin** Compute a Gram-Schmidt decomposition of $G = B^t D B$, with B orthogonal and D diagonal; $V \leftarrow BV$; **for** $(j = n; j \geq 0, j--)$ **do** $X[j] \leftarrow \lfloor W[j]/D[j, j] \rfloor$; **for** $(i = 0; i \leq n; i++)$ **do** $W[i] \leftarrow W[i] - X[j]B[j, i]D[i, i]$; **end** **end** **return** X ;**end****Algorithm 1:** Babai's nearest Plane algorithm

More formally, this gives Algorithm 1. If the input basis is of sufficient quality, we can deduce estimates on the quality of the output result. We quote the following classical result without a proof (this result is usually stated for an LLL-reduced basis, but adapts to the general case). The family $(e_i^*)_{0 \leq i \leq n}$ denotes the basis given by the Gram-Schmidt orthogonalization of the basis $(e_i)_{0 \leq i \leq n}$.

Theorem 1 *If we apply Algorithm 1 above to the FP-appr problem, we obtain a polynomial $P = \sum_{i=0}^n x_i e_i$ such that*

$$\|P - p_E(f)\|_2 \leq \frac{\gamma^{n+1}}{2\sqrt{\gamma^2 - 1}} \|\tilde{P} - p_E(f)\|_2,$$

for any $\gamma \geq \max_{0 \leq i \leq n-1} \|e_i^*\|_2 / \|e_{i+1}^*\|_2$ and \tilde{P} is the actual solution to the FP-appr problem.

A consequence is the fact that the most orthogonal the basis, the best Babai's algorithm. This suggests again to use a preprocessing step, as described above.

5.1.3 Exact method

Exact methods, which are all a variant of an algorithm initially due to Kannan [14] consist in starting with an approximate solution, i.e. a vector w in the lattice such that $R := (v - w | v - w)$ is small. Then, the goal is to find a vector x in the ellipsoid $(v - x | v - x) \leq R$. This is done by finding a larger region containing this ellipsoid, and which can be easily enumerated. Note that once a good approximation is found, it can be used to restrict again the set of points to enumerate, since we are interested in a smaller ellipsoid. We do not give further detail, and refer to [1] for pseudo-code for this method.

5.2 Short review of preprocessing

The accuracy (approximate methods) / efficiency (exact method) of those methods highly depend on the geometry of the basis of our lattice L .

For instance, the first algorithm becomes optimal if the basis is orthogonal; and the most orthogonal the basis, the best Babai's algorithm performs. Finally, if the basis is more orthogonal, the ellipsoid in the exact method is much easier to enumerate. This leads to the idea that one should try to use the most orthogonal basis possible. This suggests to use a lattice basis reduction algorithm as a preprocessing step. In this paper, we chose to use LLL-reduced bases (see [15], computable in polynomial time) or Korkine-Zolotareff reduced bases (see [12], reasonably efficient for small dimensions); one of the reasons for this is the fact that software for computing those bases is widely available, eg. in NTL [19].

It should be pointed, however, that our experiments showed that, in the case of polynomial approximation, the bases are sufficiently orthogonal, and in practice Babai's algorithm always gave the optimal solution. One might, however, expect that the situation changes if the dimension increases too much (say, above 50), but this does not seem to be the case for applications.

5.3 Choice of the precision

An important question when looking for an approximation is the choice of the functions e_i . Indeed, if one is looking for polynomial, or trigonometric approximations, one shall work with $x^i/2^{m_i}$ or $\cos(ix)/2^{m_i}$. Choosing all the m_i equal to zero would yield a very poor approximation ! We shall thus look for e_i under the form $E_i/2^{m_i}$, where E_i are the "actual functions" by which we want to approximate, eg. $E_i = x^i$, or $E_i = \cos(ix)$ and try to find an optimal choice for the m_i .

Indeed, the choice of the m_i is of direct consequence on the quality of the final approximation, but this is not the sole factor. Indeed, the error $\|P - f\|_2$ can be decomposed as the sum of two factors, one coming from the projection of f over E , namely $\|f - p_E(f)\|_2$, and the error in approximating $p_E(f)$ by an element of our lattice. Increasing the m_i only affects this second term, whereas increasing n affects mostly the first term. It thus appears pointless, once n is fixed, to increase the m_i beyond reason. We give two heuristics on the choice of the m_i .

Heuristic 1 *Let G be the Gram matrix $(E_i|E_j)$. If P is the solution to our CVP problem, we expect that*

$$-\log_2 \|P - p_E(f)\|_2 \approx \frac{\sum_{i=0}^n m_i}{n+1} - \frac{\log_2 \det(G)}{2(n+1)}.$$

In particular, it is pointless to choose the m_i much larger, on average, than $-\log_2 d_E(f)$.

A first possibility is to choose all the m_i to be $\lceil -\log_2 d_E(f) \rceil + \alpha$ for a small value of α . We can however try to estimate which of the m_i should be made larger because the corresponding vector carries "more information".

Heuristic 2 *m_i should be chosen roughly equal to*

$$\alpha - \log_2 d_E(f) + \log_2 \|E_i\|_2.$$

Indeed, if the basis E_i were orthogonal, and if $p_E(f) = \sum_{i=0}^n \varphi_i E_i$, the error can be written as

$$\|P - p_E(f)\|_2^2 = \sum_{i=0}^n \left| \frac{p_i}{2^{m_i}} - \varphi_i \right|^2 \|E_i\|_2^2.$$

The i -th term in this sum is expected to be of the order of $2^{-2m_i-2} \|E_i\|_2^2$. Thus, m_i should be taken as $\log_2 \|E_i\|_2 + C$ in order to minimize the global error. Since for an orthogonal basis, $2 \sum_{i=0}^n \log_2 \|E_i\|_2 = \log_2 \det G$, we deduce Heuristic 2 from Heuristic 1.

In practice, experiments show that the basis is “sufficiently” orthogonal for the heuristic above to give results close to optimal.

5.4 Numerical issues

The main numerical issues of our algorithm are

- the computation of orthogonal bases which is required at almost every step of the method,
- the problem of computing accurately enough the integrals that define the matrix G ,
- the definition of the problem itself, namely the fact that we shall have to replace exact real numbers (the value of the integral) by floating-point (or, in practice, integer) approximations.

We shall take the second item for granted, eg. by using [7]. The first item is a classical topic and has been extensively studied, see for instance [16]. We shall however focus on the last point. The meaning of this is that we replace the exact values of G and V by approximations \tilde{G} and \tilde{V} . A first (trivial) remark is that the problem is already ill-defined in dimension 1 : finding the element of $\mathbb{Z}\pi$ closest to $\pi/2$ is highly subject to numerical approximation. However, the *distance* to $\mathbb{Z}\pi$ is well-defined: the solution found might change, but we are switching from an optimal solution to another solution which is arbitrarily close of being optimal. In the example above, depending on the roundings, we might find either 0 or π , both being admissible answers to the original problem.

Theorem 2 *For M a matrix, we define $\|M\|$ to be the maximum of absolute values of its coefficients, and similarly for a vector. Assume that $\max(\|G - \tilde{G}\|, \|V - \tilde{V}\|) \leq \varepsilon$, and that $n\varepsilon \|\tilde{G}^{-1}\| < 1$. Then, for all $x \in \mathbb{Z}^n$, we have*

$$\left| (x - G^{-1}V)^t G (x - G^{-1}V) - (x - \tilde{G}^{-1}\tilde{V})^t \tilde{G} (x - \tilde{G}^{-1}\tilde{V}) \right| \leq \varepsilon c(x, n, \tilde{G}, \tilde{V}),$$

where $c(x, n, \tilde{G}, \tilde{V})$ is an explicit constant depending only on $x, n, \tilde{G}, \tilde{V}$.

Proof. We have

$$(x - G^{-1}V)^t G (x - G^{-1}V) - (x - \tilde{G}^{-1}\tilde{V})^t \tilde{G} (x - \tilde{G}^{-1}\tilde{V}) = V^t G^{-1} V - \tilde{V}^t \tilde{G}^{-1} \tilde{V} - 2x^t (GV - \tilde{G}\tilde{V}).$$

The last term is at most, in absolute value

$$\begin{aligned} n\|x\| \|(G - \tilde{G})V - \tilde{G}(V - \tilde{V})\| &\leq n^2\varepsilon\|x\|(\|V\| + \|\tilde{G}\|) \\ &\leq n^2\varepsilon\|x\|(\|\tilde{V}\| + \|\tilde{G}\| + \varepsilon). \end{aligned}$$

The difference $V^t G^{-1} V - \tilde{V}^t \tilde{G}^{-1} \tilde{V}$ can be rewritten as $(V - \tilde{V})G^{-1}V + \tilde{V}(G^{-1} - \tilde{G}^{-1})V + \tilde{V}\tilde{G}^{-1}(V - \tilde{V})$ whose norm is bounded from above by $n^2\varepsilon\|G^{-1}\| + n^2\|G^{-1} - \tilde{G}^{-1}\|\|\tilde{V}\|(\|V\| + \varepsilon) + n^2\varepsilon\|\tilde{G}^{-1}\|\|\tilde{V}\|$.

Write $G - \tilde{G} = E$ with $\|E\| \leq \varepsilon$, i.e., $G = \tilde{G}(I_n + \tilde{G}^{-1}E)$. Under our assumptions, the series $\sum_{n \geq 1} (-1)^n (\tilde{G}^{-1}E)^n$ is convergent, and thus

$$G^{-1} = \left(I_n + \sum_{n \geq 1} (-1)^n (\tilde{G}^{-1}E)^n \right) \tilde{G}^{-1},$$

which proves that

$$\|G^{-1} - \tilde{G}^{-1}\| \leq \frac{n\varepsilon\|\tilde{G}^{-1}\|^2}{1 - n\varepsilon\|\tilde{G}^{-1}\|},$$

which concludes the proof of the theorem. \square

The constant can be easily deduced from the proof by putting all error terms together.

This theorem shows that a “good” vector for the approximate problem is not too far away from a “good” vector for the exact problem, and (by symmetry) vice-versa.

The drawback is that the quality of the approximation depends on x . However, since x lies in a bounded region of \mathbb{R}^n , one can in theory remove this dependency, or check once x is computed that the precision is sufficient. Using the theorem (and the theorem where we exchanged the roles of G , V and \tilde{G} , \tilde{V}) shows that by increasing the precision we can come as close as we want to the actual optimal.

6 Some experiments, comparison with L^∞ -approximation

We have implemented the various combinations of methods described above and tested them for the problem described above on several functions. The algorithms have been implemented in C, using NTL-5.4[19], GMP-4.2.1 [8] and MPFR-2.2.0[18]. The scalar products have been computed using GP/pari function `intnum`, but we plan to replace it with software yielding guaranteed results, such as the CRQ library [7].

These implementations of the LLL algorithm may present numerical instability [16], but this is not expected on matrices of the kind we reduce, in the dimension useful for applications. In any case, this does not compromise the optimality of the results computed; simply, the preprocessing is of lesser quality than expected, and thus the computation time slightly larger.

All these computations, even in larger dimension, have taken less than a second on a Intel Pentium M 2GHz.

Example 1 We consider the function $x \mapsto \sin(\pi\sqrt{x})/(\pi\sqrt{x})$ on the interval $[0, 1]$ and we search for a degree-8 polynomial approximation with single precision floating-point coefficients. We get an L^2 error equal to $1.883 \dots \cdot 10^{-21}$, that we compare to $2.562 \dots \cdot 10^{-17}$ which is the error given by the rounded projection i.e. the polynomial obtained after having rounded to the nearest the coefficients of the real optimal L^2 -approximation: we get a factor 10000 improvement.

We obtain an absolute error equal to $1.345 \dots \cdot 10^{-10}$, that we have to compare to $1.002 \dots \cdot 10^{-8}$ which is the error provided by the rounded minimax (i.e. the polynomial obtained after having rounded to the nearest the coefficients of the real optimal L^∞ -approximation): we obtain an improvement by a factor 100.

Example 2 We consider the arctan function on $[-1, 1]$. Then, we search for a polynomial of the form $p(x) = x + x^3(p_0 + p_1y + \dots + p_{22}y^{22})$ where $y = x^2$ and the p_i are double precision floating-point numbers. This is a practical example from [6, Chap. 10]

We immediately obtain an approximant giving rise to a relative error equal to $2.71 \dots \cdot 10^{-18}$ which improves by a factor 5 the relative error given by the rounded minimax which is equal to $1.15 \dots \cdot 10^{-17}$.

There is currently another lattice reduction based approach [3] developed for tackling the problem of getting very good L^∞ -approximation. On the few experiments we did, we noticed that this approach sometimes leads to better results than ours but the factor between the errors they reach and ours is not large, even in degrees around 25. That approach has the drawback to require as input the computation of a minimax approximation, given by Remes' algorithm, whereas our approach is entirely done at almost no cost. Moreover, there are issues that the approach of [3] is not yet able to address but which can be treated with the method developed here. For instance, we already noticed that we can tackle with relative error approximation. We are also able to compute approximations in which the values of some coefficients are fixed. Finally, we can directly adapt our method to compute approximations to two-variable functions.

7 Conclusion and further work

We have presented a set of methods for computing best (absolute or relative) L^2 floating-point approximations, either by polynomials, or more generally by any finite linear combination of functions, such as trigonometric polynomials. These methods can be seen as a further tool in the toolbox available to the “computer arithmetician” to compute approximations with floating-point (and more generally machine-number) coefficients.

We have reviewed the main features of this tool: efficiency (especially in large dimension), ability to deal with relative error, and versatility, utility as a preprocessing step to speed up L^∞ or relative error computations.

The fact that the best vector produced by the L^2 method seems in practice to be within a small factor of the L^∞ solution suggests an improved strategy. Denote by b the best L^2 -approximation that we have obtained. By using a straightforward adaptation of the optimal algorithm, we can list all vectors v for which

$$\|v - p_E(f)\|_2 \leq \beta \|b - p_E(f)\|_2, \quad (2)$$

for some constant $\beta > 1$.

Then, we can compute the corresponding L^∞ norm of all those vectors, and look for the best of them. The main trouble is the fact that the number of vectors will grow as

$$\left(\frac{\beta \|b - p_E(f)\|_2}{2(\det(G))^{1/2n}} \right)^n;$$

if the m_i are not chosen carefully enough, $\|b - p_E(f)\|_2$ may be much larger than $(\det G)^{1/2n}$.

More intuitively, $\|b - p_E(f)\|_2$ is of the order of magnitude of the size of the largest vector in a “good” basis; if we have many “small” vectors in the basis, any “not too large” linear combination of those vectors can be added to b without changing significantly its L^2 -distance to $p_E(f)$. In that case, the set of vectors which verify (2) may be very large, and should be dealt with in a non-exhaustive way.

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